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EP 1 176 976 B1

(12)

EUROPEAN PATENT SPECIFICATION

(51) Int CL:

- (45) Date of publication and mention of the grant of the patent: 12.07.2006 Builetin 2006/28
- A61K 38/24(2008.01) A61P 15/08(2008.01)
- (86) International application number: PCT/GB2000/001745

(11)

- (21) Application number: 00927534.8 (22) Date of filing: 05.05.2000
- (87) International publication number: WQ 2000/067778 (16.11,2000 Gazette 2000/46)
- (54) USE OF LH ADMINISTERED IN MID- OR LATE-FOLLICULAR PHASE FOR THE TREATMENT OF ANOVULATORY WOMEN

VERWENDUNG VON IN DER MITTLERE ODER SPÄTE FOLLIKULÄRE PHASE VERABREICHTE LIH ZUR BEHANDLUNG VON ANOVULATORISCHEN FRAUEN

UTILISATION DE LH ADMINISTRÉE DANS LA PHASE FOLLICULAIRE INTERMÉDIAIRE OU TARDIVE POUR LE TRAITEMENT DE FEMMES ANOVULATOIRES

- (84) Designated Contracting States:
 AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU
 MC NL PT SE
 Designated Extension States:
 AL LT LV MK RO SI
- (30) Priority: 07.05.1999 EP 99303574
- (43) Date of publication of application: 05.02.2002 Bulletin 2002/96
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Description

[0001] The present invention relates to the use of gonadotrophins in the treatment of anovulatory women. In particular, it relates to the use of luteinising hormone (LH) for promoting follocular development, and especially paucifollicular and monofollicular development, when inducing ovulation in anovulatory women.

- [0002] Gonadotrophins are widely used in clinical practice to treat women with WHO group II and WHO group I anovulation (World Health Organisation Technical Report 514, (1973)): Conventionally, followlogenesis is induced by administering hMG (human menopausal gonadotrophin) or u-hFSH (unnary human folicle stimulating hormone) at a dose of 75 - 150 IU/day. This dose is increased after a few days (usually five) by steps of 75 IU. It is rare to exceed 450 ILI/day. When there is at least one follicle having a mean diameter of at least 18 mm and no more than two follicles having a mean diameter of at least 16 mm, a high dose (of 5000 NU for example) of hCG (human chorionic gonadotrophin) is administered to induce ovulation. This "conventional protocol" has been used successfully for more than 20 years. It carries some risks however, mainly in patients with polycystic ovarian disease (PCOD). These risks include the occurrence of ovarian hyperstimulation syndrome (OHSS), and a relatively high incidence of multiple pregnancies (Schenker et al,
- Fertil. Steril. 35:105-123 (1981)). Although the majority of multiple pregnancies are twins, induction of ovulation contributes to one third of the high rank multiple births in the UK (Levene et al, Br. J. Obstat. Gynacol. 99:607-613 (1992)). [0003] Careful monitoring during treatment by ultrasound (US) and assessment of serum oestradiol (E2) have reduced these risks but have not been able to prevent them in all patients. These problems are directly related to the difficulty of
- obtaining the growth of a single dominant folicie leading to non-physiological multifolicular development. [0004] During the last 10 years, a new protocol has been designed (the "chronic low dose protocol") and tested in order to reduce further the incidence of the complications of gonadotrophin therapy (Seibel et al, Int. J. Fertil., 29:338-339 (1984); Buvat et al. Fertil. Sterii., 52:553-559 (1989); Hamilton-Fairley et al. Human Reprod 6:1095-1099 (1991); Sagle et al, Fertil Steril, 55:56-60 (1991); Shoham et al, Fertil, Steril, 55:1051-1056 (1991); Meldrum, Fertil Steril, 55: 1039-1040 (1991)). This protocol starts with a low dose of FSH or hMG (75 IU/day) and no dose adjustment before seven or preferably 14 days of treatment. If a dose adjustment is required, this is made by incremental steps of only 37.5 IU. In addition, each subsequent increase may only be effected after seven days of treatment at a given dose. The concept of this chronic low dose protocol is to find the threshold amount of FSH necessary to promote unifoliculogenesis. Encouraging results have been published so lar, showing that this approach reduces the mean number of preovulatory
- folicies, the average preovulatory E2 level and the size of the ovary at mid-luteal phase. [0005] However, despite the use of the chronic low dose protocol, some treatment cycles still have to be cancelled due to an over-response (e.g. where there are more than 3 folicies with a mean diameter of 16 mm or more). In addition, the multiple pregnancy rate, although clearly improved when compared to the conventional protocol, is still higher then in spontaneous conception cycles i.e. 5 - 10 % in induced ovulation as opposed to 1.5 % in spontaneous cycles. This is due to the fact that development of a single pre-ovulatory follicle is obtained in only about two thirds to three quarters
- of the induced cycles and follicles having a mean diameter of 15 mm or less are usually not considered when assessing the number of pre-ovulatory follicles on the day of hCG administration (Buvat et al, Fertil, Steril, 52:553-559 (1989); Hamilton: Fairley et al, Human Reprod 6:1095-1099 (1991)). It is however not clear whether follicles with a mean diameter of 14 to 15 mm, or even less, on the day of hCG administration, will ovulate and lead to the release of a healthy fertilisable occyte. Thus, it would be desirable to have improvements in FSH-induced follocular development treatment in which the rates of multiple pregnancy and cycle cancellation are reduced.
- [0005] Antral follicle growth is induced by FSH, Continuously throughout life and up to the menopause, some follicles enter a growth phase which is interrupted by regression and atresia before reaching the full maturity stage of preovulatory status (Hillier, Hum. Reprod., 9:181-191 (1994)). During the growth phase, any follicle could be rescued from atresia, provided that it is exposed to a sufficient concentration of FSH. The level of FSH required to prevent atresia and promote
- further growth of a follicle is called the "FSH threshold" level (Brown, Aus. NZ J. Obstet. Gynecol., 18: 47-55 (1987). The FSH threshold level varies with time and, at a given time-point, the follicles which are currently in a growth phase have different FSH threshold levels. This is the rationale on which the "chronic low dose" protocol is based. A progressive and cautious increase in the dose of FSH is used for finding the threshold level of a minimal number of lollicles, and hopefully achieving mono-ovulation.
- [0007] It is known that luteinising hormone (LH) also contributes to the phenomenon of follicle dominance and monoovulation. Indeed, although some LH is essential for cestrogen synthesis during folloulogenesis, there is evidence that excessive exposure to LH will trigger follicular atresia and suppress granulosa proliferation. Developing follicles appear thus to have finite requirements for stimulation by LH, beyond which normal follocular development ceases. This is the "LH ceiling" concept (Hillier, Hum. Reprod, 9:181-191 (1994)). It is believed that, at a given time-point, the follicles which are currently in a growth phase have different LH ceiling levels. It is suggested that the more mature folloles are more
 - resistant to the atretic action of LH than less mature follicles. [0008] Two cases of WHO group I anovulation treated by either FSH alone or hMG using a step-up protocol have been reported (Glasier et al, Journal of Endocrinology, 119 A-159 (1988)). The "FSH alone" cycle had a much larger

erway no week and an analysis of the second and the

number of majure follicles than the hMG cycle, possibly supporting a role of LH in the atresia of secondary follicles. Afterwards two comparative studies were published. In a first cross-over study in 10 hypogonadotrophic hypogonadal women, a striking difference was recorded in terms of preovulatory E₂ levels, but follocular count was not reported (Couzinet et al, J. Clin. Endocrinol. Metab. 66:552-556 (1988)). A second cross-over study in 9 hypogenadotrophic hypogenadal women reported a mean number of follicles having a mean diameter of more than 16 mm on the day of

hCG administration of 2.0 (0.7 in hMG-treated cycles and of 1.2 in FSH-treated cycles (Shoham et al, Fertil. Steril., 55: 1051-1056 (1991)). No information to available on the number of smaller follicles.

- [0009] More recently, the results of administering 150 IU hFSH (human FSH) and 75 IU r-hLH (recombinant human LH) to a single patient with unmeasurably low serum FSH, LH and cestradiol concentrations have been published (Hall et al. The Lancet, 344(8918):334-335 (1994)). Administration of r-hLH and hFSH caused E₂ levels to be raised, and the total number of follicles of 10 mm or more in diameter to be reduced, as compared to administration of hFSH alone. However, the number of large follicles remained sufficiently high to suggest an unacceptably high multiple pregnancy rate. [0010] A further study compared the effect of administering r-hLH (at a dose of either 300 IU/day or 750 IU/day) and r-hFSH to normal ovulatory women after treatment with FSH for stimulating multiple follicular development prior to
- intrauterine implantation (Sullivan et al, Journal of clinical Endocrinology and Metabolism, 84, 228-232, 1999)). The results indicate that serum E2 levels were raised in those women who received LH, although no measurements of the number end size of follicias were made and a multiple pregnancy occurred in the group receiving 750 IU/day of LH. [0011] According to the present invention, there is provided the use of LH and/or a biologically-active analogue thereof in the production of a medicament for inducting pluri- or uni folliculogenesis in anovulatory women at a daily dose in the
- range of from 100 to 1500IU, wherein the medicament is to be administered starting mid- to late folloular phase. [0012] As used herein, an "IU ratio" is the ratio of the number of IU of one component to the number of IU of another component. It is notaworthy that gonadotrophins may now be expressed in (mass/µg) instead of biological IU. In this case, a conversion factor has to be used to translate the naw value into IU. For convenience, references hereinafter to LH, FSH and hCG are intended to include biologically-active analogues thereof.
- [0013] The invantors have found that the edmnistration of LH at a dose of 100 to 1500 IU/day during mid- or late follocular phase can promote paucifollicular development, that is to say, it can reduce the number of preovulatory folloces per treatment cycle in patients undergoing follicular induction, as compared to cycles where LH is not administered at a dose of 100 to 1500 iU/day. LH administered in accordance with the invention can induce unifoliculogenesis, i.e. the development of a single preovulatory folicle. Doses in the range of from 200 to 800 IU/day, and more preferably 225 to 450 IU/day, have been found to be particularly effective. The reduction in multifolicular development can reduce the number of cycles cancelled owing to excessive folicle development, i.e. it can rescue those cycles when there are an excessive number of folicies, making the process of evulation induction more efficient. in addition, the incidence of multiple pregnancy and of OHSS can be reduced.
- 100141 The required daily dose may be administered as a single dose each day. Thus, the medicament may be packeged so as to provide only the daily dose of LH, e.g. in a unit-dose container such as a vial. However, it is possible that LH may be administered on two or more occasions during the day - provided of course that total LH administered during the day equals the daily dose - and the medicament packaged accordingly, i.e. in a multi-dose conteiner. It is also possible that LH could be administered on alternate days or at even longer intervals. Such decisions will be taken by the physician administering the medicament and will depend on parameters such as the patient's body mass index (BMI), medical history, stage of folicular development when receiving LH, metabolism, response to the treatment, the

half-life of the medicament and so on.

- [0015] Folliculogenesis will generally be induced in anovulatory women by the administration of FSH using the conventional protocol or the chronic low dose protocol described abova or an alternative protocol. LH should be administered at an appropriate stage of folicular development, i.e., the mid- to late-follicular phase. This stage may be decided by the physician administering the medicament and may depend on the regime by which ovulation is induced. By way of
- example, the appropriate stage of folloular development may be judged to have been reached when at least e single follicle reaches a mean diameter of 8 mm, or when at least one follicle has a mean diameter in the range 10-15 mm (preferably 13-14 mm), or when there are more than 3 follicles with a mean diameter in the range of from 8 to 13 mm and no larger folloles.
- [0016] The administration of LH will generally ceese when ovulation is induced by the administration of the high dose of hCG. Again, the timing of hCG administration to induce ovulation may be decided by the physician. For example, it may be when there is at least one follicle having a diameter of 18 mm or more and no more than 3, preferably 2, follicles having a diameter of 11 mm or more.
- [0017] LH is to be administered only when the required stage of follocular development has been reached. In this case, the administration of FSH can be discontinued altogether or can be continued at the same dose as before, or at a lower or higher dose, it is preferred if the administration of FSH is continued but at a lower dose than previously, the dose being lower than that of LH.
 - 100181 Thus, FSH and/or a biologically active analogue thereof may be used in combination with LH in the production

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- of the medicament. In this embodiment, the IU ratio of LH to FSH is preferably in the range of from 1.5:1 to 20:1. More preferably, the ratio is in the range of from 1.5:1 to 10:1.
- [0019] When the medicament is for administration after the appropriate stage of follicular development has been reached, the IU ratio of LH:FSH may be about 10:1. A particularly preferred daily dose for such a medicament is 375 IU of r-hLH and 37.5 IU of r-hFSH.
- [0020] There is provided the use of LH and FSH and/or biologically-active analogues thereof in the production of a medicament for inducing pauci- or uni-foliculogenesis in women at an IU ratio of LH to FSH in the range of from 1.5:1
- [0021] The invention may be modified in that LH is replaced by an equivalent dose of hCG and/or a biologically-active analogue therenf
- [0022] As used herein, an "equivalent dose" of human chorionic gonadotrophin (hCG) is calculated on the basis that 1 IU of hCG is equivatent to 5-7 IU of LH in the pharmacopaeta Van Hell bloassay (Van Hell, H, et al. Effects of human menopausal gonadotrophin preparations in different bioassay methods, Acta Endocrin., 47: 409-418, 1964). For convenience, references herein to luteinising hormone (LH) are intended to include hCG, with dosas of LH being intended to include the equivalent dose of hCG.
- [0023] LH, FSH and hCG may be obtained from natural sources, e.g. Isolated from urine, pituitary or placenta, or may be obtained using recombinant DNA technology (see WO85/01959 end Loumaye et al, Human Reprod, 11: 95-107. 1996). Biologically-active analogues thereof include peptidic analogues, non-peptidic analogues and chimeras. It is preferred if human LH and FSH are used in the present invention.
- [0024] Compounds useful in the invention may be formulated for administration by any convenient route, often in essociation with a pharmaceutically and/or veterinarily acceptable carrier. It is preferred that the compounds are formulated for parenteral edministration.
- [0025] It is preferred that the LH and FSH (when present) be administered subcutaneously, preferably into the anterior llew lenimohde
- 25 [0026] Formulations for parenteral administration will usually be sterile. Phermaceutical formulations adapted for parenteral administration include aqueous and non-aqueous sterile injection solutions which may contain anti-oxidants, buffers, bacteriostats and solutes which render the formulation isotonic with the blood of the intended recipient; aqueous and non-agueous sterile suspensions which may include suspending agents and thickening agents are also within the scope of the invention. The formulations may be presented in unit-dose or multi-dose containers, for example sealed ampoules and vials, and may be stored in a freeze-dried (tyophilised) condition requiring only the addition of the sterile liquid carrier, for example water for injections, immediately prior to use. Extemporaneous injection solutions and suspensions may be prepared from sterile powders, granules and tablets. The formulations can be administered through
 - a prefilled syringe, an euto-injector or a multidose auto-injector. [0027] Oral and other enteral formulations need not be stanle and may be presented in unit- or multi-dose form. Oral tormulations may be in the form of solids, such as powders, granules, tablets, capsules (for example herd or soft galatin capsules) or lozenges, or liquids, such as syrups or elixirs. Fillers and/or carriers may be present as appropriete, and those skilled in the art of pharmaceutical formulation will be able to provide such additional or elternative excipients as may be necessary or desirable; flavouring agents are one example. Any formulation intended for oral administration may be formulated for enterio resistance, so as to assist delivery to the small intestine by avoiding or mitigating any
 - digestion of the compound(s) as may occur in the stomach or the proximal part of the small intestine. Tablets or capsules may be enteric coated, for example by conventional procedures, Liquid formulations may be effectively rendered enteric registant by including or being co-administered with a suitable agent such as medium-chain triglycerides. [0028] Enteral compositions other than oral compositions include rectal compositions, which may be in the form of a
 - suppository. Suppositories will generally include a suppository base, such as cocoa butter. Again, particular formulations containing the active ingredient(s) may routinely be prepared by those skilled in the art of pharmaceutical formulation. (0029) The invention will now be described further in the following non-limiting examples.

Example 1

- [0030] The effect of LH when administered after FSH stimulation was examined on WHO Group II anovulatory women during a clinical study conducted according to ICH GCP (International Conference on Harmonisation - Good Clinical Practice) guidelines. The patients had the following characteristics:
 - Premenopausal; aged between 18 and 39; infertile due to ovulatory dysfunction; have had spontaneous menses, menses induced by clomiphene citrate therapy or a positive progestin-induced withdrawal bleed within the previous year; a body mass index of 35 or less (calculated as body weight in kg divided by (height x weight) in m²); euthyroid; no medical condition which may interfere with the absorption, distribution, metabolism or excretion of LH; no clinically systemic disease; no known allergy to gonadotrophin preparations; no persistent ovarian cyst of 11 mm or greater

or ovarian endometrioma (as determined by ultrasound); no previous or current hormone dependent tumour; no clinically relevant reproductive tract disease; and no active substance abuse.

[031] The patients underwent routine ovalidation induction with FSH until there were 4 or more follicles in the range of rome 1-3 mm in diameter, no larger follicles and an endomentum of 8 mm or more follicles in the range of rome 1-3 mm in diameter, no larger follicles and an endomentum of 8 mm or more folliclesses. They were then randomised into 3 blinded groups, one to receive a placebo, one to receive 225 IUMay of r-hLH and one to receive 450 IUMay of r-hLH.

[0032] Table 1 below summarises the respective groups of patients:

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Table 1

| Mean ±SD | Placebo | r-hLH 225 IU/day | r-hLH 450 IU/day |
|----------------|-------------|------------------|------------------|
| No of patients | 5 | 4 | 8 |
| Age (yrs) | 29.2 ± 5.7 | 26.8± 6.2 | 30.9± 3.9 |
| (min-max) | (23-35) | (20.35) | (25-38) |
| Weight (kg) | 62.8 ± 15.9 | 60.0±1.6 | 66.8 ± 15.4 |
| (min-max) | (47-88) | (58-62) | (48-97) |
| BMI | 24.6 ± 4.7 | 22.8 ± 1.9 | 24.7 ± 4.9 |
| (min-max) | (20-31) | (21-25) | (18-34) |

[0033] r-hLH (LHadi Ø, Serono) was used in vals containing 76 IU r-hLH and 47.75 mg of sucrose, phosphate buffer and Tween 20 in a hophilised form. LHadi is produced in genetically engineered Chinese harrister Ovary (CHO) cells in which the genes encoding that alpha and beta chains of human LH have been introduced through recombinant technology. The specific activity of UHsel is approximately 15000 IU LHing.

10034]. For dose 22.5 L. 3 visia were used. Doe visi was reconstituted in 1 mil of water and gestly aptitude, skilling are an endo operative this he arbest report. The totality of the energing publical was applied and used for processing the state of the second visit. After gentle agistion, the bits lift of the resulting publical was applied and used for reconstitution of the second visit. After gentle agistion, the bits lift of the resulting publical was applied and used for reconstitution of the limit of the form of the second visit. After gentle agistion, the totality of the resulting publical was applied and mandatality injected subditionable to the second visit of the seco

[0038] The placebo was in vials matching the r-hLH vials but containing only sucrose, phosphate buffer and Tween 20. [0036] The r-hLHplacebo treatment was condinued for 7 days unless at least one folicile reached a mean diameter of at least 18 mm and there were 3 or lever folicible having a mean diameter of 11 mm or greater. In this case, e single dose of 5000 [U.O. hLGC (Protatol & Sarrono) was given subculaneously.

ose of 3000 II of II-NC4 (**Pross et, servici) was given succusined US) was used at intervals of 1-2 days to measure (0037) Prict to and during the in-Lit-place-bo treatment, uitrasound US) was used at intervals of 1-2 days to measure the mean diameter of the foliciose (determined as the mean of the two longest perpendicular diametars) and the endometrial lithichness (assessed as the distance from the hyper-choogenic interface of the endometrium and the mytometrium).

40 to the opposite interface including the stronger midline echo (endometrial interface)). All follides with a mean diameter of 11 mm or greater were recorded. (2003) Prior to and each time an utrasound scan was carried out during the r-hLH/placebo treatment, a blood sample

Lives sixth and the resulting sentin was analysed for E₂ (cestinabil), P₄ (progestionos), U.F. FSH and androstenedome, [DR33] E₂ and P₄ were analysed using DPC Gode-count, RIA sold lysaes coated table separation, LH (serring and unless) and FSH were analysed using MAIACLONE FRIMA, and androstenedone was analysed using Diagnossic System. Laboratories method. RIA.

[0040] The results are summarised in Tables 2-4 and in Figure I of the accompanying drawings which is a graph showing the size and number of follicles on the day of hCG administration (or the last day of treatment of no hCG was administrated) for each of the patients.

50 [0041] It can be seen that the administration of LH at 225 or 450 IUday subsequent to FSH treatment resulted in more marked folicular regressions than in the administration of placebs, as suggested by patients with complete folicular regression, a smaller number of folicides on the day of hCQ administration and a reduction in folicide median size from 15 mm in the folicides on the folicides of the folicid

[0042] The efficacy of r-hLH in promoting mono-ovulation is libstrated by the emergence of a dominant folicle (as evidenced by the median size), the absence of folicular phase luteinisation and a comparatively lower P_e, level at the mid-timal chance.

Example 2

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[0043] The effect of LH and FSH administered during the late follicular phase was examined on WHO Group I anoualistry women during a clinical that conducted according to ICH GCP guidelines. The patients had the following characters.

primenopausal, aged between 18 and 36; actinical history of hypogonadotrophic hypogonadism; have alloped of restment (a say) with pulsatia G-RH2, posedotrophins or cestropen progesterons treatment therapy at least one month between the screening productly, have had a register progressioner challenge test performed outling the screening princip, had the following homonal values in a fasting blood sample (between 7 and 9.30 AM) drawn within 6 months before the restment practic.

| FSH: | < 5 miU/mi |
|------------------------------------|--------------------|
| LH: | < 1.2 mlU/cni |
| Thyroid stimulating hormone (TSH): | < 6.5 µlU/ml |
| Free T ₄ : | > 11 and < 24 pmoV |
| Tastosterone: | < 3.5 nmoVI |
| Prolactin (PRL): | < 520 mIUI; |

no clinically agrillicant absorbum finding, within 6 months pilor to sady start, in pre-seament heartackopy, in official, chemistry and utinary languagement or residue of no patiboding stagetizers or durate homes multiple in the control of the prophogola stagetizers or durate homes in the line, no, offie, it is the prophogola of the p

The study was divided into an open phase of a meximum of 28 days and a binded phase of a meximum of 7 days. 0.044) In the open phase, all potential scienced 225 UMOs of r-NH and 1125 UMOs of r-NFSH. Here was no file in 5_g level or sign of foliosized growth after 7 days, he does of r-NFSH was nised to 150 UMOs, After a further 7 days, the does of r-NFSH was mised to 152. UMby if there was no fise in 5_g level or sign of foliosized growth. further 7 days, the does of r-NFSH was mised to 262.5 UMby if there was no rise in 5_g levels or sign of folicular growth. The does of r-NH remained constant introduction the cone no hase.

[0045] When a patient had at least one folicle with a mean diameter in the range of from 10.15 mm, she entend the solicited phase, in this places, the patients were anatomised into 3 bifuld groups, one to receive a LH placebo and centinus the dose of hiPSH necelved not hele stat got the peop phase, one to neceive 255 Ulday of ri-LH and conflinus that dose of rhiPSH necelved on the last day of the post phase, and one to receive 255 Ulday of ri-LH and sertificate that dose of rhiPSH necelved on the last day of the post phase, and one to receive 255 Ulday of ri-LH and a PSH placebo. (0046) Table 5 below summarises for resceived recorder of patients.

Table 5

| Mean±SD | FSH/Placebo | r-hLH/placebo | FSH/r-hLH |
|----------------|-------------|---------------|------------|
| No of patients | 6 | 6 | 8 |
| Age (yrs) | 31.9 ± 6.2 | 31.0± 3.0 | 30.8 ± 4.6 |
| (min-max) | (21-39) | (27-34) | (25-37) |
| Weight (kg) | 70.3 ±10.0 | 51.7±4.4 | 66.9 ±15.9 |
| (min-max) | (60-88) | (46-59) | (50-89) |
| BMI | 25.2 ±2.3 | 19.8 ±1.1 | 24.6 ±4.3 |
| (min-max) | (21-28) | (19-21) | (20-30) |

[0047] r-hFSH (Gonal-F Ø, Serono) was used in ampoules containing 75 IU r-hFSH and 30 mg sucross and phosphate buffer in a lyophilised form, up to 3 of which were dissolved in 1 ml of water for injection. Matching ampoules containing only sucrose and phosphate buffer were provided for the FSH placets.

[0048] r-h,H (LHadi®, Serono) was provided and administered as in Example 1. The LH placebo was in vials matching the r-h,LH vials but containing only sucrose, phosphate buffer and Tween 20.

[0049] All injections were made subcutaneously into the anterior abdominal wall.

- [0050] The blinded phase was continued for 7 days unless at least one folicite reached a mean diameter of at least 18 mm and there were 20 or fewer folicites having a mean diameter of 11 mm or greater. In this case, a single dose of 10000 IU of th/GC (Profasi 6), Serono) was given substaneously.
- [0051] On the first, fifth and eight days of the open phase, and at regular intervals (i.e. 1 to 2 days) during the blinded phase, ultrasound was used to measure the mean diameter of the follicles and the endometrial thickness. All follicles with a mean diameter of 11 mon greater were recorded.
- [0052] On the first day of the open phase, and at regular intervals (i.e. 1 to 2 days) during the blinded phase, a blood sample was taken and the resulting serum was analysed for E₂, P₄. LIL FSH and androstenedione as in Example 1.
- (0053) The results are summarised in Tables 6-9 and in Figure 2 of the accompanying drawings which is a graph showing the size and number of folicides on the day of hCG administration (or the last day of treatment of no hCG was administration) for each of the patients.
- [0054] it can be seen that stopping FSH and administering r-hLH at 225 ItJ/day resulted in a marked and excessive folloular regression.
- [0055] The efficacy of r-hLH in promoting mono-ovulation in the presence of FSH is illustrated by a reduction in the mean number of folicides having a diameter of 14 mm or greater, an increase in the proportion of patients with only 1 or 2 folicides having a diameter of 14 mm or greater, the emergence of a dominant folicide (as evidenced by a me dan folicide size of 12 mm as compared to 15 mm for the FSH placeds group), and the absence of folicidar phase laterials of the contract of the co

| . Treatment Group | Patient Id | Number of Folicles > =8 mm at Baseline | Number of Folides > =11 mm Last US | Number of Follicles > = 14 mm Last US | hCG Received | Reason / Comment |
|----------------------|------------|---|---|--|--------------|-------------------------|
| Placebo | 20002 | 8 | 5 | 4 | No - | Risk of OHS |
| | 30003 | 22 | 14 | 10 | Yes | |
| | 40001 | 20 | 3 | 2 | Yes | |
| | P40005 | 12 | 3 | 1 | Yes | |
| | P40008 | 8 | 3 | 2 | Yes | |
| | N = 5 | 14.00±6.63 | 5.60±4.77 | 3.80±3.63 | 4 Yes/ 1 No | |
| r-hLH 225 IU/day | 20001 | 5 | 1 | 0 | No. | Follicles regressed |
| , | 30001 | 12 | 0 | 0 | No | Failure of treatment |
| / | 40003 | 18 | 5 | 2 | Yes | |
| | 40007 | 4 | 3 | 3 | Yes | |
| | N = 4 | 9.75±6.55 | 2.25±2.22 p=0.4391 | 1.25±1.50 ρ=0.2342 | 2 Yes/2 No | |

| | | | (continued |) | | |
|---------------------|------------|--|---|--|--------------|---------------------------------|
| Treatment Group | Patient Id | Number of Follicles > =8 mm at Baseline | Number of Follicles > =11 mm Last US | Number of Follicles > = 14 mm Last US | hCG Received | Reason / Comment |
| r-hLH 450 IU/day | 10001 | 6 | 0 | 0 | No | all follicles became atretic |
| | 20003 | 10 | 13 | 4 | No | Risk of OHSS |
| | 30002 | 9 | 5 | . 3 | No | Failure of treament |
| | 40002 | 17 | 3 . | 1 | Yes | |
| | | | | | 1 1/ | |

Yes

No

624.25 Yes

1.50±1:41.

Failure of treatment

4 Yes/4 No 3.75±410 p=0.8684 p=0.2731 p-values from comparison with placebo group (ANCOVA adjusted for number of follicles at baseline)

3.

P: pregnant patient

20

30

40009

50001

70001

N=8

8.88±3.83

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| | | ı | | | | | Admi | Administered | | | | |
|----------------|---------------------|---|--------|--------|----------------------|----|-----------|----------------------------------|------------|----------|------------|---------|
| | | Ц | Ę | catmen | Treatment Randomised | B | | | | a-values | lines | |
| | | Ĺ | Macebo | 7 = | IV/day | 3 | r-hLH 450 | Contrast" | One-sided | | Two-sided | 2 |
| Variable | Number of Follicles | z | P.C | z | 82 | z | 180 | | Asymptotic | React | Asymmetric | Bytes |
| Polliches > == | 0 foll.> = 1 Imm | 0 | 90.0 | E | 25.0% | 2 | 25.0% | Placebo vs. r-hLH 225 IU | 0.0562 | 0.1429 | 0 1124 | VPLI 0. |
| E I | 1 foll.>=11 mm | 0 | 20.0 | - | 25.0% | 0 | 0.03 | Placebo vs. r-hLH 450 IU | 0.1108 | 0.2319 | 0.2217 | 0 1600 |
| | 2 foil. > = 13 mm | 0 | 900 | • | 0.0% | 0 | 0.0% | F-MLH 225 IIJ vs. F-M. H 450 III | 0 7987 | 0.1870 | 1,000 | 2010 |
| | 3 foll. > = 11 mm | _ | 60.09 | - | 25.0% | 4 | \$0.0% | Overall comparison | 0.2064 | 0 2222 | 0.4178 | 0 4178 |
| | >3 foll >= 11 mm | 7 | 40.0% | - | 25.0% | 7 | 25.0% | | | | | 0.427.0 |
| | ΝV | ~ | 100.0% | * | 100.0% | 80 | 100.0% | | | | | |
| Follicles > ** | 0 foll. > = ! 4mm | 0 | 90.0 | 2 | \$0.0% | 2 | 25.0% | Placeby vs. r-hLH 225 IU | 0.0774 | 0.1429 | 0.1547 | 0.2857 |
| 14 mm | 1 foll. > = 14 mm | - | 20.0% | 0 | 0.0% | ~ | 37.5% | Placeho vs. r-hLH 450 fU | 0.0817 | 0.1298 | 0.1635 | 0.2416 |
| | 2 foll. > = 14 mm | 7 | 40.0% | - | 25.0% | - | 12.5% | C-MLH 225 IU M. F-MLH 450 IU | 0.3786 | 0.4788 | 0.7577 | 0 8171 |
| | 3 foli. > = 14 mm | • | 80.0 | 7 | 25.0% | - | 12.5% | Overall comparison | 0.1259 | 0.1354 | 0.2510 | 5292.0 |
| | >3 foll. > = 14 mm | 7 | 40.0% | 0 | 0.03 | - | 12.5% | * | | | | |
| | 411 | Ŀ | 20000 | Ŀ | 100 000 | ŀ | 1 | | | | _ | |

Overall Comparison; Jockheere-Terpstra test Pairwise Comparison: Cochran-Armitage test for trend.

Table 4 – Descriptive Statistics of Hormone Levels Measured at 71 and on the Day of bCG or on the Last Day of Treatment If No bCG was Administered

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| | | | | TI (first | T1 (first day of stimulation) | dation) | | | Day of | CG or last | Day of hCG or last day of treatment if no hCG was | mens if no | hCG was |
|-----------------|------------------|----|--------|-----------|-------------------------------|---------|-------------|-----|--------|------------|---|------------|-------------|
| Variable | Treatment | = | Mean | g | маѕ | Median | Range | a | Mean | as | SEM | Median | Rance |
| PSH (IU/L) | Placebo | ~ | 12.20 | 9.60 | 2.50 | 9.50 | (8-21) | 2 | 6.54 | 3.84 | 77.7 | 2.8 | (2-11) |
| | r-hLH 225 IU/day | 4 | 12.53 | 6.75 | 3.37 | 11.45 | (6-21) | 4 | 7.35 | 2.98 | 1.49 | 6.03 | (6-12) |
| - | r-bLH 450 IU/day | ~ | 9.1 | 3.7 | 5 . | 9.80 | (8-19) | • | 6.94 | 2.02 | 0.73 | 6.25 | (2-10) |
| | All | 91 | 11.80 | 4.84 | 1,21 | 9.63 | (6-21) | 17 | 6.92 | 2.70 | 99.0 | 6.20 | (2-12) |
| LIE (IU/L) | Placebo | s | 7.84 | 6.30 | 2.82 | 4.80 | (3-18) | \$ | 6.12 | 2.74 | 1.23 | 6.90 | (6-2) |
| | r-hLH 225 IU/day | 4 | 5.25 | 5.69 | .3 | 6.20 | (1-7) | 4 | 6.30 | 4.48 | 2.24 | 6,40 | (2-12) |
| | r-hLH 450 1U/day | 7 | 3. | 4.1 | 1.57 | 3.40 | (1-13) | , | 19.9 | 4.14 | 1.56 | 4.80 | (3-15) |
| | ΥII | 9 | 5.78 | 4.58 | 1.15 | 4.55 | (91-1) | 9 | 6.53 | 3.60 | 9.90 | 6.00 | (2-15) |
| E2 (proof/L) | Placebo | 2 | 4031.6 | 3759.9 | 1681.5 | 3612.0 | (21001-865) | \$ | 4780.6 | 4612.7 | 2062.9 | 3540.0 | (313-11040) |
| | r-hLH 225 IU/day | * | 1491.8 | 1633.5 | 816.8 | 851.5 | (384-3880) | 4 | 2550.0 | 4715.7 | 2357.8 | 227.0 | (153-9633) |
| | r-hLH 450 IU/day | 1 | 1376.7 | 885.8 | 334.8 | 1315.0 | (123-2809) | ó | 1966.9 | 2665.1 | 942,3 | 297.0 | (133-7269) |
| | ΥI | 91 | 2235.1 | 2486.8 | 621.7 | 1304.5 | (123-10017) | (1) | 2934.0 | 3763.6 | 912.8 | 378.0 | (133-11040) |
| P4 (ranol/L) | Placebo | 2 | 4.56 | 1.80 | 08'0 | 4.30 | (3-7) | 5 | 8.86 | 10.83 | 4.84 | 4.50 | (2-28) |
| | r-hLH 225 1U/day | 7 | 3.08 | 25 | 0.51 | 3.05 | ÷ | 4 | 2.68 | 1.15 | 0.57 | 2.50 | 6-6 |
| | r-hLH 450 JU/day | , | 2.47 | 1.03 | 0.39 | 2.30 | 9 | 10 | 2.89 | 1.73 | 19.0 | 2.25 | 9-1) |
| | Ail | 2 | 3.28 | 1.53 | 0.38 | 2.85 | (1-3) | 2 | 4.59 | 6.24 | 1.51 | 2.80 | (1-28) |
| Androstenedione | Placebo | 5 | 17.42 | 11.62 | 5.20 | 16.50 | (\$-32) | 2 | 15.74 | 7.03 | 3.14 | 14,40 | (8.27) |
| (nmol/L) | r-NLH 225 IU/day | 4 | 8.63 | 9.88 | 0.44 | 8.30 | (8-10) | 4 | 11.75 | 1.92 | 96.0 | 12.00 | (9-14) |
| | r-hLH 450 IU/day | • | 10.53 | 7.11 | 5.69 | 8 | (5-26) | 00 | 12.18 | 9.5 | 3.38 | 8.95 | (6-35) |
| | All | 91 | 12.21 | 8.38 | 2.09 | 01.6 | (5-32) | 11 | 13.12 | 7.49 | 1.82 | 11.40 | (36.35) |

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| | | | | | | | | | EP 1 1 | 76 9 | 76 1 | B1 | | | | | | | | | | |
|---|---------------------------|----------------------|----------------|-------|-------|-------|-------|--------------|-------------------|-------|-------|-------|-------|-------|---------------|-------------------|-------|-------|-------|-------|-------|-------|
| | hCG Received | No | o _N | No | Yes | Yes | Yes | 3 Yes/3 No | No. | Ŷ. | S. | Yes | No | Yes | 2 Yes/4 No | Yes | oN. | Yes | Yes | Yes | S. | Yes |
| | Cumulative LH Dose | | | | | | | | 929 | 1125 | 1675 | 1125 | 1576 | 006 | 1162.6±360.5 | 675 | 450 | 226 | 675 | 450 | 1575 | 875 |
| ancellation Blinded Phase | Cumulative FSH Dose | 225 | 450 | 338 | 150 | 300 | 899 | 337.3±150.0 | | | | | | | | 563 | . 225 | 113 | 338 | 522 | 788 | 450 |
| s and hCG C | Number of Days | 87 | 8 | 8 | | . 2 | 9 : | 2,7±1,4 | 8 | 2 | ۷. | 2 | . 7 | 4 | 5.2±1.6 | er - | 21 | - | ຄ | 2 | 4 | 3 |
| Table 6 - Summary Data on Stimulation Open and Blinded Phases and hCG Cancellation Onen Phase Blinded P | Cumulative LH Dose | 1800 | 2825 | 1575 | 2250 | 2925 | 1575 | 2175.0±631.1 | 3375 | 450 | 1675 | 1575 | 1125 | 2700 | 1800.0±1064.9 | 3825 | 675 | 2475 | 1675 | 1575 | 1350 | 2475 |
| imulation Open a | Cumulative FSH Dose | 006 | . 1688 | 788 | 1238 | 1688 | 788 | 1181.3±425.4 | 1988 | 525 | 788 | 788 | 563 | 1538 | 981.3±654.9 | 2363 | 338 | 1238 | 788 | 788 | 675 | 1388 |
| y Data on Stim | Numberof | 80 | 13 | 7 | 9 | 13 | 7 | 9.7±2.8 | 15 | 2 | 7 | 7 | 2 | 12 | 8.0±4.7 | 17 | 8 | Ξ | 7 | 2 | 8 | 11 |
| ole 6 - Summar | LastDose of FSH (IU) | 112.5 | 150.0 | 112.5 | 150.0 | 150.0 | 112.5 | 131.3±20.5 | 150.0 | 112.5 | 112.5 | 112.5 | 112.5 | 150.0 | 125.0±19.4 | 187.5 | 112.6 | 112.5 | 112.5 | 112.5 | 112.5 | 150.0 |
| Tak | First Dose of FSH (IU) | 112.5 | 112,6 | 112.5 | 112.5 | 112.6 | 112.6 | 112.5 | 112.5 | 112.5 | 112.5 | 112,5 | 112.5 | 112.5 | 112.5 | 112.5 | 112.5 | 112.5 | 112.5 | 112.5 | 112.5 | 112.5 |
| | Patient Id | 10002 | 10004 | 20001 | 30005 | 40002 | 50001 | 9=2 | 10003 | 10005 | 30003 | 40001 | 50002 | 60002 | N=6 | 10001 | 10006 | 20002 | 40003 | 40004 | 50003 | 50004 |
| | Treatment Group | Gonal- F/ Placebo | | | | | | | r-hLH/ Placebo | | | | | | | Gonal-F/r- hLH | | | | | | |

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| | _ | | _ | $\overline{}$ | |
|------------|---------------|---|-------|------------------------------------|--|
| , | | hcG Received . | oN . | 5 Yes/3 No | |
| 0 | | Cumulalive LH Dose | 450 | 365.6±223.0 646.9±406.7 5 Yes/3 No | |
| 5 | Blinded Phase | Cumulative FSH Dose | 225 | 365.6±223.0 | |
| 0 | | Number of Days | 2 | 2.9±1.8 | |
| e Q | | Cumulative LH Number of Cumulative Cumulative LH Dose Dose PSH Dose Dose | 1350 | | |
| (confined) | | Cumulative FSH Dose | 675 | 8.5±4.3 1031,3±632.0 1912,5±977.1 | |
| 5 | Open Phase | Number of Days | 9 | 8.5±4.3 | |
| 0 | | Treatment Patient Id First Dose Last Dose of Number of Group FSH (IU) FSH (IU) Days | 112.5 | 126±27.9 | |
| 15 | | First Dose of FSH (IU) | 112.5 | 112.5 | |
| 10 | | Patient Id | 10009 | N=8 | |
| i6 | | Treatment Group | | | |
| | | | | | |

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Table 7 - Summary Data on Number and Size of Follicles and hCG Cancellation

| - 1 | | | ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, | Last US | | | |
|-----|---------------------|------------|---|----------------------------------|-----------------------------------|-----------------|---|
| | Treatment Group | Patient Id | Number of Foliclés > = 10mm on T1 | Number of Follicles > = 11 mm | Number of Folicles >= 14 mm | hCG Received | Reason/ Comment |
| , | Goual-F/ Placebo | 10002 | 1 | 5 | 1 | No | possiblerisk of multiple pregnancy |
| | | 10004 | 2 | 4 | 2 | No | 1 follicle 18 mm plus 3> 11 mm; not within protocol |
| | | 20001 | 1 | 4 | 3 | No | multiple follicles |
| | | 30002 | 4 | 5 | 4 | Yes | |
| ' | | 40002 | 1 | . 3 | 2 | Yes | |
| | | 50001 | 2 | 4 | 4 | Yes | |
| 5 | | N=6 | 1.83±1.17 | 4.17±0.75 'p=0.5008 | 2.67±1.21 *p=0.4071 | 3 Yes/3 No | |
| | r-hLH/ Placebo | 10003 | 1 | 0 | 0 | No | regression of follicles |
| , | | 10005 | 1 | 1 | 0 | No | regression of follicles |
| | | 30003 | 2 | 0 | 0 | No | failure of treatment |
| | | 40001 | 2 | 3 | 2 | Yes | |
| 5 | | 50002 | 2 | 4 | 0 | No | fallure of treatment |
| | | 60002 | 1 | 1 | 1 | Yes | |
| | | N=6 | 1.50±0.55 | 1.50±1.64 **p=0.0171 | 0.50±0.84 **p=0.0162 | 2 Yes/4 No | |
| | Gonal-F/r- hLH | 10001 | 1 | 4 | 2 | Yes | |
| | | 10006 | 1 | 13 | 1 | No | risk of OHSS |
| | | 20002 | 2 | 2 | 2 | Yes | |
| | | P40003 | 4 | 3 | 1 | Yes | |
| | | P40004 | 3 | 3 | 1 | Yes | |
| , | | 50003 | 2 | 0 | 0 | No | failure of treatment |
| | | 50004 | 1 | 4 | 1 | Yes | |
| 5 | | 60001 | 2 | 19 | 8 | No | risk of OHSS |

-- - --- ---

.....

| | | | Last US | | | |
|--------------------|------------|--|----------------------------------|------------------------------------|-----------------|--------------------|
| Treatment Group | Patient Id | Number of Follicles > = 10mm on T1 | Number of Folficles > = 11 mm | Number of Folicies > = 14 mm | hCG Received | Reason/ Comment |
| | N=8 | 2.00±1.07 | 6.00±6.50***p=0.0032 | 2.00±2.51 ***p=0.0412 | 5 Yes/3 No | |

p values adjusted for BMI: contrast p-value with the previous treatment group

^{*} Gonal-Fir-hLH vs. Gonal-FiPlacebo ** Gonal-FiPlacebo vs.r-hLHIPlacebo

^{***} r-hLH/Placebo vs. Gonal-Fir-hLH

P: pregnant patient

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| | | | | | | | | 9/ | 01 | • | | | | | |
|----------------------|---------------|--------------------|---------------------|----------------------------------|-----------------------------------|--------------------------------|--------------------|---------------------|---------|----------------------------------|-----------------------------------|--------------------------------|--------------------|--------------------|--------|
| | | page | Fraci | 0.0281 | 0.2887 | 0.1016 | 0.4325 | | | 0.0152 | 0.1485 | 0.1575 | 0.2769 | | |
| | 20 | Two-sided | Asymptotic | 0.0115 | 0.1641 | 0.0799 | 0.4101 | | | 0.0092 | 0.0868 | 0.0922 | 0.2660 | | |
| | p-values | 9 | Exact | 0.0141 | 0.1538 | 0.0633 | 0.2184 | | | 0.0076 | 0.0765 | 0.0276 | 0.1377 | | |
| | i | One-sided | Asymptotic | 0.0057 | 0.0820 | 0.0399 | 0.2051 | | _ | 0.0046 | 0.0424 | 0.0461 | 0.1330 | | |
| | | Contract | | Gonal-F/Placebo vs r-hLH/placebo | Gonal-F/Placebo vs Gonal-P/r-hL.H | 1-hLH/Placebo vs Gonaf-F/r-hLH | Overall comparison | | | Gonal-F/Placebo vs r-hLH/placebo | Gonal-P/Placebo vs Gonal-P/r-hL.H | r-hLH/Placebo vs Gonol-Fir-hLH | Overall comparison | | |
| | | Gonal-F r- | % | 12.5% | %00 | 12.5% | 25.0% | \$0.0% | 100.0% | 12.5% | 50.0% | 25.0% | 0.0% | 12.5% | 100.0% |
| *** | | <u>.</u> | z | Ŀ | • | _ | ~ | 4 | * | Ŀ | • | ^ | • | = | - |
| Transment Dandamicad | THE PARTY AND | r-bLH Placebo | × | 33.3% | 33.3% | 0.0% | 16.7% | 16.7% | 100.0% | 66.7% | 16.7% | 16.7% | 0.0% | 0.0% | %0'001 |
| 1 | | _ | z | 7 | ~ | ۰ | = | - | 9 | | - | _ | ٥ | ۰ | 9 |
| ľ | | Gonal-F Placebo | % | %000 | 2000 | 0.0% | 16.7% | 83.3% | %0'001· | %00 | 16.7% | 33.3% | 16.7% | 33.3% | 100.0% |
| L | | | z | ۰ | • | ۰ | - | ~ | 9 | 0 | - | ~ | - | 7 | ٥ |
| | | | Number of Follicles | 0 (oll. > e.) Imm | I foll. > + 11 mm | 2 foll. > = 11 mm | 3 foll.> = 11 mm | > 3 foll. > = 11 mm | ΝY | 0 foll. > = 14mm | 1 fall. > = 14 mm | 2 foll. > = [4 nm | 3 foll. > = 14 mm | >3 foll. > = 14 mm | l V |
| | | | Variable | Follicles | mm! !** | | | | | Follicles | >= 4mm | | | | |

Overall Comparison: Jonekheere-Terpsira iesi Pairwise Comparison: Cochran-Arminge iest for tread

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| | | | | TI (first d | TI (first day of stimulation) | (uoiseio | | L | Day of hi | Day of hCG or last day of treatment if no hCG was | sy of treatn | rent If no ht | 3G was |
|-----------------|-----------------|----|---------|-------------|-------------------------------|----------|------------|----|-----------|---|--------------|---------------|------------|
| | | | | | - | | | | | ä | administered | | |
| Variable | Treatment | = | Mean | SD | SEM | Median | Range | - | Mean | gs | SEM | Median | Range |
| FSH (IU/L) | Gonal-F/Placebo | 9 | 8.58 | 3.19 | 1.30 | 8.03 | (5-[4) | 'n | 8.52 | 3.13 | 1.40 | 7.00 | (5-12) |
| | r-hLH/Flacebo | ý | 12.37 | 6.73 | 2.72 | 53.6 | (9-36) | 9 | 3.33 | 2.13 | 0.87 | 3.00 | (9-1) |
| | Goral-F/r-hLH | 80 | 9.68 | 3.44 | 1.22 | 10.15 | (4-15) | 80 | 9.03 | 5.66 | 25 | 9.55 | (4-13) |
| | Ali | 20 | 10.16 | 4.62 | 1.03 | 9.75 | (4-26) | 61 | 7.09 | 3.62 | 0.83 | 7.00 | (1-13) |
| LH (IU/L) | Gonal-F/Placebo | 9 | 80'1 | 91.0 | 20'0 | 0.1 | (1-1) | n | 8: | 00:0 | 0.00 | 1.00 | <u>:</u> |
| | r-hLH/Phocebo | 9 | 9 | 99.0 | 0.24 | 8 | 3 | 9 | 1.88 | 1.56 | 0.64 | 1.25 | 5 |
| | Gonal-F/r-hLH | 80 | 1.58 | 0.87 | 0.31 | 1.10 | (4-3) | 8 | 1.56 | 0.71 | 0.23 | 1.35 | (6-1) |
| | Ail | 50 | 1.35 | 0.65 | 0.15 | 1.00 | (6-1) | 16 | 1.52 | 87 | 0.23 | 8 | (5:0) |
| E2 (pmol/L) | Gonal-F/Placebo | 9 | 05'169 | 737.24 | 300.98 | 474.5 | (160-2171) | 2 | 725.80 | 99.686 | 442.59 | 302.00 | (163-2483) |
| | r-hLHVPlacebo | 9 | 669.33 | 483.03 | 197.20 | 630.00 | (129-1311) | ۰ | 116.33 | 102.12 | 41.69 | 100.00 | (33-316) |
| | Gonal-F/r-hLH | 00 | 1416.50 | 1666.01 | \$89.02 | 650.00 | (187-4885) | - | 3452.86 | 3843.18 | 1452,59 | 1537.00 | (251-1257) |
| | AI. | 20 | 974.85 | 1167.89 | 261.15 | 474.50 | (129-4885) | 81 | 1583,17 | 2803.84 | 650.87 | 309.00 | (33-11257) |
| P4 (nmol/L) | Gonal-F/Placebo | 9 | 4.1 | 9:0 | 0.2 | 3 | (1-1) | ~ | 9.1 | 8.0 | 0.4 | 1.3 | (1-3) |
| | r-hLH/Placebo | 9 | 7.7 | 2 | 6.0 | 9: | 2 | 9 | 6. | 2 | 9.0 | 1.2 | 5 |
| | Gonal-F/r-hLH | 8 | 2.3 | 1.3 | 0.5 | 2.0 | (1.5) | 7 | 22.9 | 53.0 | 20.0 | 2.8 | (2-143) |
| | A | 20 | 2.0 | = | 0.2 | 9.1 | (1.5) | 18 | 0.01 | 33.2 | 7.8 | 1.9 | (1-143) |
| Androstenedione | Goral-F/Placebo | 9 | 4.87 | 2.55 | 1.04 | 3.65 | (3-10) | 7 | 4.08 | 2.35 | 2.1 | 3.20 | (2.8) |
| (mod/L) | r-hLH/Placebo | ø | 5.93 | 2.50 | 25 | 3.65 | 66 | 9 | 5.63 | 2,37 | 0.97 | 5.13 | 6-10 |
| | Gonal-F/r-hLH | 8 | 7.71 | 3.78 | 1.34 | 7.30 | (3-14) | • | 10.58 | 6,19 | 2,19 | 10.80 | (3-22) |
| | 7 | 8 | 6.33 | 3.19 | 0.71 | 5.85 | (3-14) | 8 | 7.48 | 5.18 | 1.22 | 06 \$ | 12.22) |

Claims

15

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- The use of LH and/or a biologically-active analogue thereof in the production of a medicament for inducing paucifoilibulgeness or unfolliculogenesis in anovulatory women at a daily dose in the range of from 100 to 1500 IU, wherein the medicament is to be administered starting in the mid-lo- late-folicular phase.
- The use as claimed in claim 1, wherein the medicament is to be administered when there are more than 3 follows with a mean diameter in the range of from 8 to 13 mm and no larger follows.
- 3. The use as claimed in claim 1 or claim 2, wherein the LH is r-hLH.
- 4. The use as claimed in claim 1, 2 or 3, wherein the daily dose is in the range of from 200 to 800 IU.
- 5. The use as claimed in claim 4, wherein the daily dose is in the range of from 225 to 450 IU.
- The use as claimed in any preceding claim, wherein FSH and/or a biologically-active analogue thereof is used in the production of the medicament.
- 7. The use as claimed in claim 6, wherein the IU ratio of LH to FSH is in the range of from 1.5:1 to 20:1.
- The use as claimed in claim 7, wherein the ratio is in the range of from 1.5:1 to 10:1.
- The use as claimed in any preceding claim, modified in that LH and/or a biologically-active analogue thereof is replaced by an equivalent dose of hCG and/or a biologically-active analogue thereof.

Patentansprüche

- Verwendung von LH und/oder einem bologisch-aktiven Analogon devon, in der Herstellung von einem Medikament zum Induzieren der Paut-Dillitutogenese oder Unifolizuitogenese in annovalstorischen Frauen bei einer täglichen Desie im Bereich von 100 bis 1500 IU, wobei das Medikament mit Beginn der mittleren bis späten folliktidiren Phase verabreicht wird.
- Verwendung nach Anspruch 1, wobei das Medikament zu verabreichen ist, wenn mehr als 3 Foliikel mit einem mittleren Durchmesser im Bereich von 8 bis 13 mm und keine größeren Foliikel vorhanden sind.
- 3. Verwendung nach Anspruch 1 oder Anspruch 2, wobei das LH r-hLH ist.
- 4. Verwendung nach einem der Ansprüche 1, 2 oder 3, wobei die tägliche Dosis im Bereich von 200 bis 800 IU ist.
- Verwendung nach Anspruch 4, wobei die t\u00e4gliche Dosis im Bereich von 225 bis 450 IU ist.
- Verwendung nach einem der vorangehenden Ansprüche, wobel FSH und/oder einem biologisch-aktiven Analogon davon in der Herstellung von einem Medikament verwendet wird.
- 7. Verwendung nach Anspruch 6, wobei das IU-Verhällnis von LH zu FSH im Bereich von 1,5:1 bis 20:1 ist.
- 8. Verwendung nach Anspruch 7, wobei das Verhältnis im Bereich von 1,5:1 bis 10:1 lst.
- Verwendung nach einem der vorangehenden Ansprüche, so modifiziert, dass LH und/oder ein biologisch-aktives Analogon davon durch eine äquivalente Dosis an hCG und/oder einem biologisch-aktiven Anelogon davon ersetzt ist.

Revendications

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 Utilisation de la LH et/ou d'un anaiogue actil sur le plan biologique de celle ci dans la production d'un médiciament pour induire une paucioliticulognetes ou une unifoliculognetes chez une ferme à cycle a novulatoire à une dose quoticienne dans intervalle de 100 à 1500 UI, dans laquelle le médicament et destiné à une administration com-

mençant au milieu de la phase folliculaire ou tardivement dans calle-d.

- 2. Utilisation suivent la revendication 1, dans laquelle le médicament est destiné à une administration lorsque sont présents plus de 3 foillicules présentant un diamètre moyen dans l'intervalle de 8 à 13 mm et pas de foillicules plus grands.
- 3. Utilisation suivant la revendication 1 ou la revendication 2, dans laquelle la LH est la r-hLH.
- 4. Utilisation suivant la revendication 1, 2 ou 3, dans laquelle la dose quotidienne se situe dans l'intervalle de 200 à
- 5. Utilisation suivant la revendication 4, dans laquelle la dose quotidienne se situe dans l'intervalle de 225 à 450 UI.
- 6. Utilisation suivant l'une quelconque des revendications précédentes, dans laquelle on utilise la FSH et/ou un analogue actif sur le plan biologique de celle-d dans la production du médicament.
- 7. Utilisation suivant la revendication 6, dans laquelle le rapport en UI de la LH à la FSH se situe dans l'intervalle de 1,5:1 à 20:1.
- 8. Utilisation suivant la revendication 7, dans laquelle le rapport se situe dans l'intervalle de 1,5:1 à 10:1.

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9. Utilisation suivant l'une quelconque des revendications précédentes, modifiée en ce que la LH et/ou un analogue actif sur le plan biologique de celle-ci est remplacé par une dose équivalente d'hCG et/ou d'un analogue actif sur te plan blologique de celle-cl.

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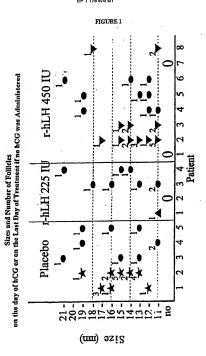
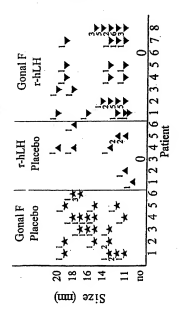


FIGURE 2



on the Day of hCG or the Last US if no hCG was Administered

Individual Size and Number of Follicles